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# Particulars of MONAI (Laid-Open) Specific

patent KOKAI (Dald-Open) No. 100760 (83)

laid-Open Data: June 18, 1983

Request for Examination: None

patent Applicataion No. 200861/81

Application Date: December 15, 1981

Convention Priority(ies): None

Inventor(s)

Applicant: Asahi Dow Dtd.

Title of Invention:

lower-temperature heat-shrinkable multilayered barrier film and a process for preparing the same

### SPECIFICATION

1. Title of the Invention:

Lower-temperature heat shrinkable multilayered barrier film and a process for preparing the same

- 2. Claims: ;
- (1) An oil-resistant lower-temperature heatshrinkable multilayered barrier film which is a four or
  more-layered film comprising at least one layer of a blend
  composition (ABC) layer consisting of

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(A) at least one polymer selected from copolymers of ethylene and monomers selected from vinyl-ester monomers,

- (1), wherein said polymer (A) is a polymer which is a appolymer of ethylene and at least one monomer selected from the group consisting of acrylic acid, acrylic acid esters, methacrylic acid and methacrylic acid esters and has a content of said monomer of 1 to 12 moles.
- (1), wherein said polymer (A) consists of the linear low-density polyethylene and has a melt index of 0.2 to 10 and a density of 0.910 to 0.935 g/cm<sup>3</sup>.
- (5) The multilayered barrier film according to claim (1) or (4), wherein said polymer (A) is the linear-low-density polyethylene prepared by copolymeriting ethylene with 7 moles or less of at least one clefin selected from a-clefins of 3 to 12 carbon atoms such as propylene, butene, pentene, hexene, heptene, octene, 4-methyl-l-pentene and the like as the  $\alpha$ -olefin.
- (5) The multilayered barrier film according to claim (1), wherein said polymer (Ε) is a copolymer of 93 mole% or less—and 40 mole% or more of ethylene and said α-olefin is selected from propylene, butene-1, 4-methyl-1-pentene and the like.
- (1) or (6), wherein said polymer (3) is a soft copolymer which is prepared by randomly copolymerizing ethylene with an a-clefin and a small amount of polymes and has a Vicat softening point of 80°C or below.
- (8) The multilayered barrier film according to claim

heat-shrinkable multilayered barrier film comprising respectively melt kneading a blend composition (ASC) consisting of

- (A) at least one polymer selected from copolymers of athylene and monomers selected from vinyl ester monomers, aliphatic unsaturated monocarboxylic acids and alkyl esters of said monocarboxylic acids and linear low-density polyethylene,
- (B) a soft elastomer, consisting of an ethylene-a-clefin copolymer and having a density of 0.91  $\rm g/cm^3$  or below,
- either of crystalline polypropylene, crystalline polybutene-1 or a mixture thereof, a vinylidene chloride copolymer (PVD) having the crystal melting peak within the range of 140 to 155 °C measured by using a differential scanning calcrimetry (DSC) and a resin (S) consisting of a polymer consisting mainly of linear low-density polyethylene (LLDPE) selected from the polymer (A), providing at least one (ABC) layer adjacent to the (FVD) layer, providing at least one (S) layer on the surface layer, extruding the resulting 4 or more-layered tubular raw film through a multilavered die, quenching and solidifying the raw film with a liquid refrigerant, directly or heating the prepared film to 100°C or below and cold drawing said film at a drawing temperature within the range of 30 to 90°C at a surface area draw ratio of 4 or above and 30 or less, thereby to afford a drawn film comprising the PVD

film surface during drawing in the direumferential direction, stretching and drawing the film and deflating the drawn film simultaneously with the maximum diameter attained.

### 3. Detailed Explanation of the Invention:

This invention relates to a four or morelayered lower-temperature heat-shrinkable multilayered parrier film comprising a vinylidene chloride copolymer as a core layer. More particularly, it relates to a novel heat-shrinkable film having excellent various characteristics, such as lower-temperature heat shrinkage characteristics (shrinkage factor and shrinkagae stress), low-temperature toughness, high strength of sealed parts (oil resistance, heat resistance and impact resistance at low temperatures of sealed parts), gas barrier properties, optical characteristics, such as transparency after heat shrinking, ply separation resistance, airtightness in clipping and the like. Uses thereof are not limited, and the film is usable for skin packaging, nonshrink packaging, flexible deep drawing packaging and the like in addition to shrink packaging.

As an example of uses thereof, the shrink packaging is taken and detailed hereinafter.

The presence of heat-shrinkable films for tightly shrink packaging contents is hitherto known for long. There have been various studies aiming at exhibi-

DTXs: backsdrud what pe calling one at a pray spankade factor and stress with not only lower-temperature shrinkability in parts contacting packaged materials but also at a sufficiently lower temperature in parts out of contact therewith in order to obtain sight packages including the above-mentioned contents and the like. However, if the surinkaje characteristics are expidiced to a high tempersoure side or temperature dependence of shoinkage factor is great, i.e. in the case of films having shrinkage cheracteristics of rapidly shrinking at a certain camperature, phenomena of discoloring or datariorating contents of uneven shrunkeye due to temperature difference are caused, directly resulting in creases, sagging of the lika. Thereby defects are often developed in that the surfaces of packages are hard to see or breakage will occus therein during transportation or the like.

Therefore, shrinkable films are required to have dimensional stability at a film preservation temperatures and exhibit shrinkage characteristics to the side of lower temperatures in shrink packaging and small temperature dependence. On the other hand, gas barrier properties are characteristics required for enhancing preservation quality of contents for a long particl, and shrink package ing of particularly facty foods cannot be considered in the absence of the characteristics.

Rowever, there is nothing if films satisfying the above-mentioned two characteristics are opaque,

content, for example is and law by weight, and improved adhagive properties to PVD copolymer resin on both sides of the PVD copolymer containing 7 to 10% by weight of a plasticiter on the basis of weight to form three layers. drawing the resulting film and making the large amount of the plasticizer migrate from the layer consisting of the FVD copolymer, thereby improving barrier characteristics or the like. In such films, barrier characteristics are unstable with time and simultaneously deficient both in heat and oil resistance. Furthermore, there are problems, such as tendency to stickiness of films, insufficient modulus of elasticity or the like. Films are prepared by irradiating and crosslinking one side of such EVA (side) sealed on the inside when formed into a bag) with electron rays in order to improve processing stability [Japanese Patent Application KOKAI (Laid-Open) No. 34565/72]. Such processes have disadvantages in being costly and complicated, difficult sealing of sealed parts rather than improving oil resistance by crosslinking and deteriorating high-temperature resistance and high-temperature oil resistance or the like.

As a method for partially improving the above-mentioned problems, Japanese Patent Application KCKAI (Laid-Open) Nos. 47079/77, 148577/77, 82888/78, 89945/81 and the like are known as methods for providing an EVA layer or low-density polyethylene layer on one side of a

inventor has finally completed this invention.

Thus, this invention consists as follows:

- "Arfour or more-layered film comprising at least one layer of a blend composition (ASC) layer consisting of
- (A) at least one polymer selected from copolymers of ethylene and monomers selected from vinyl ester monomers aliphatic unsaturated monocarboxylic acids and alkyl esters of said monocarboxylic acids and linear lowdensity, polyethylene,
  - (5) a soft elastomer, consisting of an ethylener a-clefin copolymer and having a density of 0.91  $\rm g/cm^3$  or
  - (C) at least one polymer selected from crystalline 5e10W1 polypropylene and crystalline polybutene-1, said one layer being provided adjacent to a barrier layer of specific PVD, the surface layer of said film being provided with at least one resin (S) layer which consists mainly of linear low-density polyethylene (LLDPE) selected from said polymer (A) and has the total thickness of 2 to 25 y at a ratio of 5 to 40% to the total layer thickness. The process for preparing the film comprises highly stretching the respective layers at sufficiently lowertemperatures, i.e. below the crystal melting points of the resins which are a principal and secondary components of the aforesaid blend composition, preferably below the Vicat softening point (hereinafter referred to as VSP) of the bland composition.

in miscibility and extrudability of the ray material. If the melt index is higher than 10, strength as a base gaterial may be insufficient. For example, a bubble undesirably tends to readily break in stretching or the like. Among them, EVA is most preferred for use as the blend composition layer, and the vinyl acetate group content thereof is preferably 3 to 3 moles, more preferably ly 3 to 7 mole3. The linear low-density polyethylene (LIDPE) refers to linear low-density polyathylene obtained by a medium-, low-pressure or, in some cases, highpressure process, and prepared by copolymerizing 7 moles or less, preferably about 1 to 5 moles of especially at least one clefin selected from d-clefins of 3 to 12 carbon atoms, such as propylene, buttene, pentene, hexene, heptene, octene, 4-methyl-l-pentene or the like as an a-olefin. The melt index thereof is preferably 0.2 to 10, and the density is preferably 0.910 to 0.935  $g/cm^3$ . The crystal melting temperature (mp) thereof obtained by a differential scanning calorimetry (DSC) (measured at a scanning speed of 10°C/min) is 110°C or above and up to 125°C and distinguished from the crystal melting temper ature of 100 to 108 °C of branched low-density polyethylene, having a density of 0.915 to 0.927 c/cm and prepared by an ordinary high-pressure process.

The thermoplastic elastomer (E) consisting of an ethylene-c-clefin copolymer refers to a soft copolymer of ethylene and one or more clefins selected from c-clefins

Analt index of 0.1 to 10, preferably 0.2 to 6 is exempled in the form of pluried. Such a copolymer is supplied in the form of pluries without causing cold flow, as opposed to blocks in the case of ordinary unvulcanized rubber. The copolymer the case of ordinary unvulcanized rubber, the copolymer preferably has sufficient thermoplasticity so that, for preferably has sufficient thermoplasticity so that, for example even a simple substance, can be extrusion processed into the form of a film.

Inhe polymer (C) is now prystalline polypropylene and high-molecular weight prystalline polyputane-1 (hareinafter respectively abbreviated to IFF and PB-1) consisting of relatively rigid components having a relatively high degree of prystallinity. Such polymers preferably consist of relatively rigid polymers having a vicat softening point of 100 °C or above. IFF which is one of the polymer (C) refers to ordinary commercially available polymer (C) refers to ordinary commercially available polypropylene having high isotacticity, and preferably includes propylene homopolymer, or copolymers of propylene and 7 moles or less of a-olefins, such as ethylene, butene-1 or the like. Such respective copolymers may be optionally mixed.

The melt flow index is 0.1 to 30, preferably 0.5 to 20, more preferably 0.7 to 15. If the melt index is below the above-mentioned values, problems are caused in miscibility, optical characteristics and the like in processing. If the melt flow index is above the aforemaid values, problems are caused in extrusion stability.

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And the like of films are deteriorated. If the amounts the too large, films have tendencies to too much softential blocking, deteriorating heat resistance, sealing and ing, blocking and the like.

The component (C) has effects on synergistic on lesson in tensile, and impact strength, heat resistance, extrusion moldability, modulus of elasticity, and heat sealing range with other components of the blend composition, particularly remarkable effects on heat and oil resistance, extrusion moldability, modulus of elasticity, heat sealing range, partial bearing of force for preventing migrag whitening phenomena from occurring in the adjacent PVD layer in use, particularly in high shrinking and the like. The effects are as follows: If the blended amounts are small, for example processability of films and uneven section by flow characterics in a die are deteriorated and the like and deficient in heat sealing range and heat resistance of the sealed parts. In particular, oil resistance of sealed parts are insufficient at high temperatures. Conversely, if the blended amounts are too large, low-temperature shrinkability, extrusion moldability, transparency, flexibility, impact resistance and the like are deteriorated and the like. Therefore, the blended amounts are preferably within the Aforesaid range. The component (A) consists of preferably Aspecific ethylenic copolymer among those described above, and is sometimes a principal component in the blend and film temperature oil resistance of sealed parts and film grength or the like when formed into films. The lower in the density is due to the process for preparing Me resin, and values beyond the upper limit tend to no increase unstability to stretcharility as in the case of the aforesaid upper limit of the melt index and deterioration in optical characteristics of cold drawn films, particularly those after shrinking and the like (for example, Haze value glossiness or the like). It is clear that such various factors are conversely remarkably improved within the above-mentioned range by synergistic effects with other layers without deteriorating the Eforementioned processability and various characteristics and impairing various characteristics of the other layers, aspecially the ABC layer. In particular, various strength, sealing and high-temperature oil-resistance pharacteristics are markedly improved. The peak value of the crystal melting temperature (mp) measured by the DSC method at a heating speed of 10°c/min) is preferably 110 to 125 °C.

The linear-low-density polyethylene which is the principal component may be mixed with other polymers for isse so that the aforesaid various characteristics are not itself damaged, and the limits thereof are about 50% by reight or less of other components to be mixed.

One object of combination of the above-mentioned ayer construction is to synergistically improve various

large amount (for example, 7 to 12% by weight) is undesirable for improving stretchability or etrudability of the PVD. One reason therefor is as follows: Stretchability is hard to sufficiently carry out usually by a well-known method unless a large amount of plasticiters is used. In the process of this invention, devices are considered in selection of specific compositions and combination thereof even for layers other than the aforesaid pvD layer without requiring the use of the plasticiters. In this process, the aforementioned plasticiters used in a large amount sometimes destabilize stretchability and converely cause puncture and the like. As another reason, disadvantages are cited as follows: The above-mentioned plasticizers used in a large amount greatly deteriorate cxygen-barrier performance of films, resulting in deficient high barrier performance. Attempts have been well known for providing a layer readily absorbing the aforesaid plasticizers, for example an EVA layer with a The second commence of the second is the second sec high vinyl acetate group content (VAc) of 28% by weight adjacent thereto, absorbing the plasticizers in aging for a long period and even slightly improving the barrier properties in order to prevent the disadvantages. In such processes, quality is unstable and different from that of this invention.

As a further reason, ply separation is extremely readily caused by using a large amount of the plasticizers, and various characteristics, such as heat

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and have influence as improved performance of cold resistance, various strength, lower-temperture shrinkage characteristics and optical characteristics before and after shrinking.

Moreover, preferably, non-migratory highmolecular plasticipers such as EVA, oligomers, rubber-like substances which do neither make the appearance opaque nor cause ply-separation can be used. In that case, the addition amount of said substances is preferably 1 to 15 wt.%, more preferably 2 to 10 wt.%.

pyD having a relatively low polymerication degree and good fluidity may be further used as a base. In this case, carbonization phenomena by decomposition tend to reduce and improve processing stability in extrusion and stratchability.

Such a kind of PVD is usually, particularly in an unstretched state, brittle and broken after stabiliration even if quenched and processed into a film form, and has been considered undesirable. If the PVD is stretched at a higher temperature of about 84 to 105°C, as in the above-mentioned Japanese Patent Application KOKAI (Laid-Open) Nos. which are conventional examples, than AND THE WASHINGTON THE PROPERTY OF A PROPERT this invention, poor films result. The PVD is of the type suitable for sufficiently cold stretchability to exhibit synergistic effects with specific other layers. Thus, unplasticized films free of a substantially low-molecular migratory plasticizer can be also obtained.

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uses requiring the oxygen barrier properties so much or conversely cases where presence of the barrier properties has a difficulty.

The upper limit of the above-mentioned thickness constitution ratio is a limit required to keep lower-temperature shrinkage characteristics or cold resistance, sealability, other various characteristics, particularly low-temperature characteristics.

The upper limit of the thickness thereof may be such that the upper limit of the aforesaid thickness constitution ratio is kept, and a too thick layer is practically unnecessary for practical use.

The (S) layer forming the surface layer has the sum of thickness at a ratio of preferably 5 to 40% based on the total thickness, and the range of the thickness is preferably 2 to 25  $\mu$ , more preferably 3 to 20  $\mu$ . The lower limit thereof is a thickness required to provide a role as the surface layer.

The upper limit is set, since function of the ABC layer is deteriorated if the thickness is too great. If the resinthas low stretchability as the surface layer under the conditions of this invention, the ratio thereof is preferably low. The blend resin layer consisting of the ABC constituting the principal component has a thickness ratio of 25 to 90% except the surface layer in 65 to 95% exclusive of the PVD layer in principle.

The whole thickness is normally 30 to 100  $\mu_{\rm c}$ 

markedly deteriorating merchandise value but also resulting in ply separation and deterioration in strength of
sealed parts. The phenomena were hardly noted by shrinking at 85°C in all the cases where a PVD layer having an
ing within the above-mentioned range is used to constitute
a specific ASC layer and all the layers are stretched at a
sufficiently low temperature under the conditions of this
invention. If PVD having an mp over the upper limit of
layer is used, a tendency of tigtag whitening to occur is
observed even if the ASC layer is provided. The cold
resistance has also a tendency to deteriorate. As
described above, this tendency is noted even if a large
amount of a plasticizer (6% by weight or more) is used in
the PVD layer.

The PVD layer having an mp below the aforesaid lower limit has a tendency to deteriorate stretching stability, barrier properties, heat resistance and the like.

As mentioned above, the film of this invention is capable of exhibiting unprecedented improved physical properties by synergistic effects of the specific layer of a specified PVD copolymer with layers on both sides thereof.

As one of the greatest characteristics indicating lower-temperature shrinkability thereof, the film of this invention has heat shrinkage factor values of at least 15 and 25%, preferably 20 and 30%, more preferably

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value thereof at a lower temperature provide great advantages in that the film which is a packaging material can be packaged without disadvantages. i.e. deterioration of films (because of melting and discrienting), and deteriorating various characteristics (strength, strength of sealed parts, optical characteristics and the like) and further remarkable effects on prevention of deterioration in quality by producing broth (drip) and by boiling in case a material to be packaged is raw meat or the like.

well balanced both characteristics provide excellend creaseless packages.

One of other characteristics is improved cold-resistant impact strength, and the film has a falling weight impact strength of 150 kg·cm or above, preferably 170 kg·cm or above measured at 5°C according to the ASTM D1709-75. Commercially available products (a) and (b) respectively have values of 130 kg·cm and 145 kg·cm: but some may have a value of 230 kg·cm in Example described below. In the values, the film of this invention is excellent due to sufficient cold stretching of all the layers and impartment of high orientation. It is found that the FVD layer is most inferior in cold resistance and the film is broken therefrom to propagate the breakage and cause burst. The film of this invention is considered at a high level due to high orientation by synergistic effects of the whole layer. Since the strength is too high, measurement is hard by an ordinary method, and the following

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at a temperature of 95°C, and especially improved as compared with a value of 3 and 20 g/5 mm width of commercially available products (a) and (b) poor in oil resistance. This is an important factor which means that packaging can be safely carried out while preventing occurrence of drip (broth) due to tight shrinkage in vacuum packaging oily food, particularly oily processed meat, such as ham, bacon, sausage or the like, then shrinking the film and improving appearance or simultanecusly sterilizing the food or the like. This is evidence indicating that the film of this invention is motorer typeput begreintergiel felbriefenken produkter i del begreinster begreinter begreinte begreinter begreinter begreinter begreinter begreinter begreinte begreinter begreinte begreinter begreinte begrei especially improved in such various characteristics. Referring to Fig. 3, advantageousness of this invention is clearer.

Excellent optical characteristics before and after shrinking may be cited as one of the greatest features of this invention. This is excellent in not only optical characteristics in the initial shrinking but also high shrinkage factor after shrinking. In short, for example the rate of change in Haze value is generally small for shrinkage factor. This value is 15% or less, preferably 10% or less expressed in terms of Haze value after 10% shrinkage. More important is practically a value after high shrinking. Such parts are often present in edge, free, clip, sealed parts, crease parts or the like in actual packaging, naturally causing parts of a high shrinkage factor.

layers, simple polymer, higher (DSC) mp of the FVD layer. lower orientation degree (lower stretching degree, higher stratching temperature, factors due to resin or the like), easier occurrence of peeling phenomena between the PVD layer and the adjacent layer at high temperatures. It is a matter of course that, besides the above-mentioned optical characteristics, sealing strength, cil-resistant strength, ply separation phenomena and various strengths are markedly reduced if the phenomena occur.

On closer examination, it was clear that the THE PROPERTY OF THE PROPERTY O commercially available film (a) caused zigzag whitening phenomena at a shrinkage factor of 43% at 90°C, and the commercially available film (b) caused the whitening phenomena rapidly at 85°C after a shrinkage factor of 40%. Microscopic examination showed appearance of the phenomena here and there even before that. In comparison, no such phenomenon occurred in Example Run No. 1 of the film of this invention. All the afore-mentioned Haze values are expressed in terms of 60  $\mu$ . If the film is thicker than 60 μ, or, if necessary, colored, fabricated, embossed or laminated, the same does not apply to the case.

The film of this invention has a high tensile strength (measured by the ASTM D882-67) and normally a breaking strength of 5 kg/mm $^2$  or above, and preferably a strength of  $7 \text{ kg/mm}^2$  or above.

As one use, the film of this invention is employed by sealing in the form of a bag under severe

quenched properties and them inflated into a bubble form at a temperature of 90°C or below, preferably 35 to 80°C, more preferably 35 to 70°C and most preferably at a temperature lower than the melting point of the crystal components which are principal components of respective compositions of the above-mentioned layers, most preferably below the Vicat softening points of the original polymers which are principal components or blend under a sufficient internal pressure, for example 100 to 3000 mm H<sub>2</sub>O. Thereby a desired film is only obtained in good condition. The optimum area stretch ratio at this time varies with the respective compositions, layer constitutions, temperatures or the like, but generally 5 to 20 times, preferably 7 to 15 times, and the stretch ratio in the transverse direction carried out in a preferred case is generally 2 to 6 times, preferably 2 to 4 times. It is particularly important that conditions for providing sufficient cold stretching with prevented puncture are the respective combinataions and layer combinations within the aforesaid range at this time, and preparation of a sufficient uniform is important at the same time.

The stretching degree is as follows:

extractor established constitues and an extraction of the stability of the

After determining the stretching ratio in the longitudinal direction by the speed ratio of feed nip rolls and take-off nip rolls, the best method for most stably stretching is to seal air in a bubble, stretch the bubble to the vicinity of the stretching end point (just

position, since the raw film is particularly just under brittle temperature condition and punctured. In particular, if different kinds of resins are combined in many layers, respective stretching optimum temperatures of the resins are different, and there are many combinations THE THE PARTICULAR PROPERTY OF THE PROPERTY OF THE PROPERTY OF THE PARTY OF THE PAR incapable of stretching the whole layers. After all, impartment of orientation by drwing of any layer is often sacrificed.

Successful stretching herein described has not hitherto been achieved in the whole layers at a cryogenic temperature, for example 47°C, as in the following Examples of this invention. The stretching can only be attained by synergistic effects of using, for example a multilayered tube containing the specific aforesaid copolymer layer and uniform quenched raw film, satisfying conditions, such as a specific stretching method and the like.

Further, the heating temperature herein is the maximum temperature for the raw film before stretching. The stretching temperature herein described is a temperature of parts where the stretching is started. The temperature is further reduced to a region where the stretching ends by cooling. In the region where the stretching ends (region where the bubble reaches the maximum diameter), sufficient cooling is carried out to provide at least 40 °C or below, preferably 30°C or below, more preferably 25°C or below. Therefore, the temperature difference between the stretching THE WASHINGTON OF THE PROPERTY OF THE PROPERTY

copolymerizing 4% by weight of ethylene) and used for the blend composition (Viest softening point: 67°C) ABC... layer. LLDPE (with an MI of 2.0, a density of 0.915 g/cm<sup>3</sup>, an mp of 116°C having a peak at 120°C and a Vsp of 98°C prepared by copolymerizing 3.5 mole% of octene-1 as the a-clefin) was used as the resin (S.) for the surface layer. Furthermore, 100 parts by weight of a winylidene chloride copolymer (with a DSC peak temperature mp of 145°C prepared by copolymerizing vinyl chloride), was blended with 2 parts by weight of TVA with a VAc content of 40% by weight, 1 part by weight of epoxidized soybean oil and 0.5 part by weight of acetyl tributyl citrate to provide a PVD composition (PVD $_1$ ), which was used as a resin for the PVD layer. The above-mentioned blend compositions were respectively and separately thermoplasticized by three extruders, fused in a three-kind five-layer die, extruded at an average resin temperature of 190°C and quenched with cold water at about 8°C at a point of 5 cm from the tip of the aforementioned die to prepare tubular raw films having the respactive layers having a folding width of 120 mm and a thickness of 650  $\mu$ with uniform thickness accuracy.

The layer constitution was adjusted to provide the raw films of  $S_1 = 60~\mu/ABC_{111} = 130~\mu/PVD_1 = .100$   $\mu/ABC_{111} = 300~\mu/S_1 = 60~\mu$  from the cutside of the tube.

Such raw films were passed between 2 parts of feed nip rolls and take-off nip rolls at a higher speed

(the outermost layer was counted as the first layer hereinafter).

mable 1

Tab.	.e -				· ,
Run No.	2	3	4	5	é
constitution of product  cf product  first layer (4)  second layer (4)  maind layer (4)  fourth layer (4)	3 4 5 14 3	5 C B 21 K	6 15 0 1 3 8	6 15 15 29 15	12 10 33
Total thickness (µ)	30	50	70	80	60
Run No.  Thickness constitution of product  First layer (a)  Second layer (a)  Third layer (a)  Fourth layer (a)  Total thickness (a)  Comparative Run  and a raw film having the the raw film constitution and a raw film constitution and a raw film having the first layer. Stretching first layer. Stretching puncture in the initial tube and blowing up the entirely stretched to a raw of the resulting films available films which	tota rati same was	o of l	so µ/3' succe	of 650 0	sultir i= int

Table 2 shows values of various characteristics resulting films and two kinds of commercially available films which were comparative examples.

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William Cont. (a)	* 1 Zigzac	- ©	. 0		<b>.</b> ×	. * 2 UBG	(w1t	form	heml	uren	10 01 <b>97</b> 78
The difference of the same	Comparative	(a)		11.0 80 X	10 24 2;2 120	85	6.0	230	27	43	2.0
(cont. a)	Comparative	<b>d</b>	. 1	5.1 40 X	3 13 2.8 85	9.5	6.5	240	25	4.8	3
Table 2		<b>9</b> :		4.5	30 42 1.2	99	1.1	250	7.0	20	200

# \* 1 Zigzag whitening phenomenon

- ) i Mo occurrence at all
- Partial occurrence at a high shrinkag factor (50% or above)
- A : Occurrence on the whole surface at a high shrinkage factor (50% or above
- X : Occurrence on the whole surface at a medium shrinkage factor (about 40%)
- \* 2 Use of an edged special missile

  (with 8 grooves having a radius of 2 mm
  formed in the direction of the tip in
  hemispherical missile)

Measurement was made by cutting shrunk samples of size 10 cm x 10 cm dipped in hot water at 50°C fo 5 seconds and observing the cross section.

within a narrow range of 93 to 95°C with a partially Roccurring whitening phenomenon. Although the film of Comparative Run No. (b) was finished relatively tight at gs to 90°C, the whitening phenomenon also occurred in partially much shrunk area. Appearance of the packages was best for Run No. 1, Comparative Run Nos. (a) and (b) in the decreasing order. When such packages were cooled to 0°C and dropped from a heightlof 1 m, the film of Run No. 1 did not break the bag-even by dropping 10 times. The film of Comparative Run No. (a) broke the bag by eropping once, but that of Comparative Run No. (b) broke the bag by dropping three times. Observation after preservation at 0 to 5°C for 1 month showed that the film of Run No. 1 had both excellent appearance and quality with the least occurrence of broth (drip). This was considered as effects that shrinking could be carried out at lower temperature without leaving creases and the like on the surface. The return of color was best on opening the seal. When commercial ham was halved into blocks weighing about 1.3 kg and vacuum packaged (marginal Childinension of about 15%) tight with the respective films and dipped in hot water at 93 °C floating land, the film of (Comparative Run No. (a) peeled the surface layer part in F. about 3 seconds, and the sealed part was instantaneously Sproken and torm. The surface layer part of Comparative FRun No. (b) was dissolved and peeld in about 10 seconds, and the sealed part was broken. Such a phenomenon did not

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Thicknessess constitution	٦ .	<b>u</b> .	6	10		:
First layer (µ)»	52	en en	e G	S <sub>10</sub>	8 <sub>9</sub>	Ī
Second layer (µ)	ABC <sub>112</sub> +C <sub>1</sub>	ABC <sub>111-1</sub>	ABC <sub>112</sub>	ABC 211	ABC <sub>112</sub> 4C <sub>1</sub>	
Third layer (µ)	PVD <sub>1</sub>	PVD <sub>2</sub>	PVD <sub>2</sub>	PVD <sub>2</sub>	PVD 2	·
Fourth layer (µ)	ABC <sub>112</sub> +C <sub>1</sub> 29	ABC <sub>211</sub> +S <sub>2</sub>	ABC <sub>112</sub>	ABC211	ABC <sub>112</sub> (C <sub>1</sub>	
Fifth layer (µ)		s <sub>3</sub>	S. A.	S <sub>1.0</sub>		
Total thickness (μ)	09	62	61	6.4	59	,

```
R
Rinds of resins
  c s_{\gamma}—linear-low-density polyethylene (MI: 1.0) density:
    0.917 g/cm<sup>3</sup>, mp: 118°C, Vsp: 102°C)
  & sy-linear low-density polyethylene (MI: 05.5) density:
    lets 10 g/cm<sup>3</sup>, mp: having the peak at 118 100.0; Vsp:
    99°C)
  b Sy-linear low-density polyethylene (MI: 2.0, density:
   0.924 g/cm<sup>3</sup>, mp: 121°C, Vsp: 111°C)
  é sy-linear-low-densityh polyethylene (MI: 25, density:
    0.919 g/cm<sup>3</sup>, mp: 131°C, Vsp: 95°C)
 o sgalinear low-density polyethylene (MI: 6, density:
    0.935 g/cm<sup>3</sup>, mp: 124°C, Vsp: 118°C)
 o s7-high-pessure process low-density polyethylene
    (conventional LDPE undesignated as linear)
    (MI: 2.0, density: 0.919 g/cm<sup>3</sup>, mp: 105°C)
   o sg-medium, low-pressure high-density polyethylene
    (ordinary HDPE)
    (MI: 1.0, density: 0.950 g/cm<sup>3</sup>, mp: 132°C)
   c ABC 211-blend of (a2) 65 wt.% of EVA (VAc: 4.1 mole%, ~
     MI: 1.0, mp: 95°C, Vicat: 78°) with
     (b<sub>1</sub>) 20 wt<sup>1</sup>% of ethylene-c-olefin elastomer (as
 c described above) and
    ' (C,) 15 wt.% of IPP
    with a Vsp of 64°C
    c ABC 111-1-blend of (a1) 55 wt.% of EVA with
   (b) (b) 15 wt.% of ethylene-z-clefin elastomer and
    To(C.) 30 wt.% of IPP
```

and stable stretching could not be attained. Evne when stretching temperature was reduced to 40°C or increased to 95°C, streaks tended to longitudinally occur, and sufficient stretching could not be carried out.

Samples of Comparative Run No. 3 have a tendency similar to that of Comparative Run No. 2 and were harder to draw and readily pusture.

Samples of Comparative Aun No. 4 seemed to bubble up just before stratching though unstable, but tended to puncture, leaving streaky uneven thickness. This tendency resulted in whole whitening and opacity by increasing temperature in the same manner as above, and puncture tended to occur. Sufficient stratching could not be carried out.

Samples of Comparative Run No. 5 were instantly punctured when air was introduced into the bubble, and stretching could not be conducted at a temperature between 30 to 95°C.

Stretching of samples of Comparative Run No. 6 could be stably and continuously carried on at a sufficiently low temperature of 42°C.

Stretching was carried out by the same method.

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Stretching was carried out by the same method. that copolymers having respective mp values of 141, 149 Find 154°C expressed in terms of DSC peaks in the vinylidene chloride copolymer layers were taken as Run Nos. 12, 13 and 14 and those having mp values of 135 and 160°C were designated as Comparative Run Mos. 7 and 8. The samples of the copolymers having the mp of 141, 149 and 154 °C could be successfully and stably stretched, but the sample of the copolymer having the mp of 135°C was unstable without stopping elongation and tended to puncture in stretching. The sample of the copolymer having the mp of 160°C was hard to sufficiently and completely stretch, brittle and tended to puncture. Partial samples of Comparative Run No. 8 were simultaneously evaluated to find that optical characteristics of Run Nos. 12, 13 and 14 were much the same as those of Run No. 1. Samples of Comparative Run No. 8, however, caused marked , zigtag phenomena even with a low shrinkage factor. The Sample of Combarative Run No. 8 had insufficient shrinkage ifactors of 10% at 70°C and 15% at 80°C, a low stress of 60 = 2, a breaking strength of 4.3 kg/mm<sup>2</sup> and a falling lyeight impact strength of 40 kg cm at a low level using the above-mentioned edged missile at -30°C though heat strinkage characteristics of others were good. On the other hand, the others exhibited excellent values. This

Table 5

•							
Run No.	15 16 17	16	17	18	61	2.0	
Add ( ) (ve							<u> </u>
High polymer plasticizer	4	2	2	7	0	2	!
		,		•	_		
Acetyl tributyl citrate	c c	o	- C	: <b>-</b>	· ε	-	
Dibutyl sebacato	) ~I	: <b>-</b> -	6.0	5. 5.	ر. د 	s. o	
soybean off	٠			!		:	
Total amount of 11quid additive	-		1.5	2.5	-	5.5	

Entra de la companya No phenomenon particularly inferior to those of Page No. 1 in Example 1 was found in Run Nos. 15 to 11. and Edharacteristic values were at good levels with little difference. In contrast to this, samples of Comparative Run Nos. 9 and 10 were readily punctured, but stretching could be relatively smoothly conducted. Raw films of Comparative Run Nos. 11 and 12 were especially easily 数 blocked and unstable, and good stretching could not be practiced. Since films of Comparative Run Nos. 9 and 10 ised a special blend composition having oil resistance in the used a special blend composition having oil resistance in the use of the composition having oil resistance in the composition have been also as the composition of the composition have been also been also been also be composition of the composition of the composition have been also been also be composition of the c the layers adjacent to the PVDC layer, ply separation tended to occur in the lapse of a while after stretching. Films of Comparative Run Nos. 11 and 12 were sticky and readily blocked without stiffness. Changes of barrier properties with time were examined to obtain the following Fresults: Films of Run Nos. 15 to 20 hardly changed at room temperature with time even after 15 days and had a value of about 23 cc/m<sup>2</sup>·day·atm (23°C). Films of Comparative Run Nos. 9 and 10 showed a value of 150 cc imediately after the stretching and resulted in 120 cc. Pilasiof Comparative Run Nos. 11 and 12 exhibited a value Constitution. Some stretching and reduced to 90 and solutions and reduced to 90 and solutions. 195 Corboth at low levels. Rigger whitening occurred in News 1991 1992 of Comparative Run Nos. 9, 10, 11 and 12 by shrinking with time, especially markedly in Comparative RunNos. 11 and 12. At the time of shrinking, those of Comparative Run Nos. 11 and 12 were softened and became

WALL BOARD OF THE STATE OF THE							
3	Comparative	1 8	F	PVD 2	0 T	- 's	20
	Compar 15		. в. 3.0	PVD <sub>2</sub>	E 2		09
Table 6	Conparative 14	s <sub>1</sub>	ABC <sub>111</sub>	PVD <sub>2</sub>	ABC <sub>111</sub>	$s_1$	09
	Comparative 13	ABC <sub>111</sub>	a <sub>3</sub>	PVD <sub>2</sub>	A.)	Λ <sup>II</sup> C <sub>111</sub>	0.9
	Thickness Constitution	First layer or (p)	Second layer (H)	Third layer (µ)	Fourth layer (μ)	Fifth layer (#)	Total thickness (µ)

: :

- EVA (VAC: 10.3 moles, MI: 4, mp: 73°C, Vsp:

Note:

temperature to 95 and 100°C, deterioration in optical characteristics and reduction in lower-temperature shrinkability tended to be strong. There was a tendency to deterioration in stability of the bubble. When the temperature was further increased, only a locally thin and white elongated film of nonuniform thickness was obtained. The film of Comparative Run No. 16 was punctured and could not be drawn even at respective temperatures.

4. Brief Explanation of Drawings:

Fig. 1 illustrates temperature dependence of shrinkage factor for films of this invention and commercially available films (a) and (b), wherein

1-1: Film of this invention (Run No. 1)

1-2: The above-mentioned commercially available film (a)

1-3: The afore-mentioned commercially available
film (b).

Fig. 2 shows dependence of dart impact strength (with the aforesaid grooved edge adopted) for films of this invention, samples of Comparative example and (comparative example examp

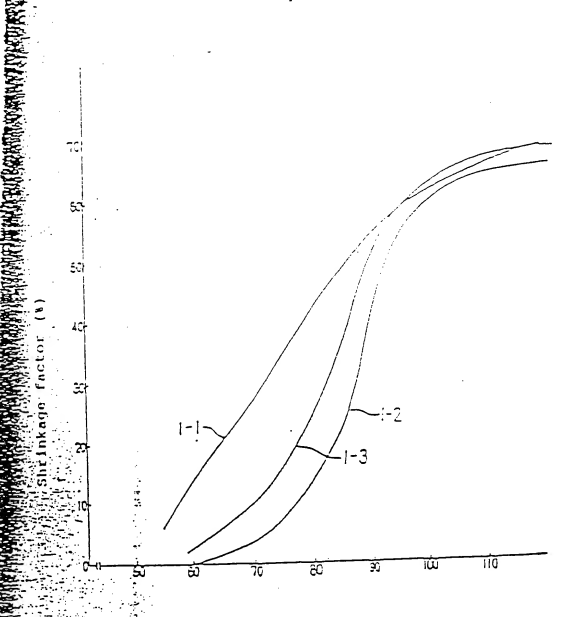
wherein

2-1: Film of this invention (Run No. 1)

2-2: Comparative Run No. 5

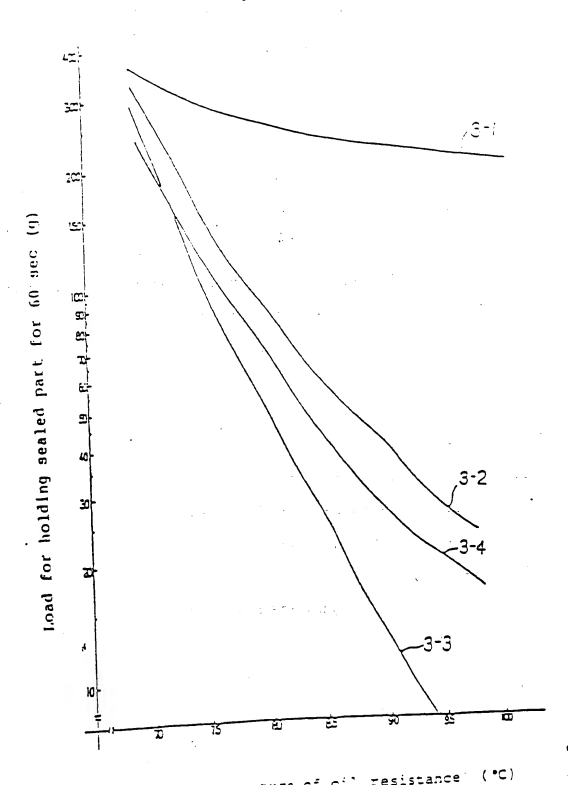
2-3: The above-mentioned commercially available





Treating temperature (°C)





Wassing temperature of oil resistance ( )